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Platypus *TCR*µ provides insight into the origins and evolution of a uniquely mammalian TCR locus1

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Abstract

TCR μ is an unconventional TCR that was first discovered in marsupials and appears to be absent from placental mammals and non-mammals. Here we show that TCR μ is also present in the duckbill platypus, an egg-laying monotreme, consistent with TCR μ being ancient and present in the last common ancestor of all extant mammals. As in marsupials, platypus TCR μ is expressed in a form containing double V domains. These V domains more closely resemble antibody V than that of conventional TCR. Platypus TCR μ differs from its marsupial homologue by requiring two rounds of somatic DNA recombination to assemble both V exons and has a genomic organization resembling the likely ancestral form of the receptor genes. These results demonstrate that the ancestors of placental mammals would have had TCR μ but it has been lost from this lineage.

Introduction

Conventional T cells exist in two distinct lineages based on the composition of their TCR heteroduplex: $\alpha\beta$ T cells use a TCR composed of α and β chains while $\gamma\delta$ T cells use γ and δ chains. Like Ig, the Ag binding V domains of the TCR chains are encoded by exons that are assembled from gene segments by somatic DNA recombination. All jawed vertebrates have both $\alpha\beta$ and $\gamma\delta$ T cells and the genes encoding these four TCR chains are highly conserved both in sequence and organization (1-3). Recently, a fifth locus encoding TCR chains, named $TCR\mu$, were found in marsupial mammals (4). $TCR\mu$ contains C regions related to TCR δ but is transcribed in a form that would include double V domains that are more related to IgH V (VH) than to TCR V genes (2, 4, 5). TCR μ does not substitute for TCR δ in marsupials since the genes encoding conventional TCR δ chains are highly conserved and expressed (2, 6).

 $TCR\mu$ genes are distinct and unlinked to those that encode conventional TCR chains and have atypical gene organization. The N-terminal V of TCR μ (V μ) is encoded by somatically recombined genes (V, D, and J), with the recombination taking place in thymocytes, resulting in clonal diversity (4). The second, C-proximal V domain (V μ j) is encoded by an exon where the V, D, and J genes are already pre-joined in the germ-line DNA and are relatively invariant (4). This is the only known example of germ-line joined V genes being used in a TCR. The $TCR\mu$ locus is also organized in tandem clusters, which is also atypical of TCR genes (2, 4).

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Searching the available placental mammal, avian, and amphibian genomes failed to uncover TCR μ orthologues (2). However, here we show that TCR μ is present in a monotreme, the duckbill platypus *Ornithorhyncus anatinus*. The monotremes are oviparous mammals that last shared a common ancestor with marsupials and placentals at least 165 million years ago (MYA) (7). The genomic organization of the platypus $TCR\mu$ locus reveals insight into the evolution of this uniquely mammalian TCR locus and supports its ancient presence in mammals.

Materials and Methods

Whole genome analysis and annotation

Analyses were performed using the platypus genome assembly Version 5.0.1 available at GenBank (http://www.ncbi.nlm.nih.gov/genome/guide/platypus/). Marsupial C μ sequences were used to search based on homology using the BLAST algorithm (4, 5, 8). Scaffolds containing C μ sequences were retrieved and exon boundaries were determined by the presence of canonical mRNA splice sites. Platypus cDNA sequences were used to search against the $\it O.$ anatinus genome project to identify the genomic V, D and J gene segments. The beginning and end of each coding exon of V, D and J gene segments were identified by the presence of mRNA splice sites or flanking recombination signal sequences (RSS). Supplementary Fig. 1 shows the location of each TCR μ V, D, J and C segments on the scaffolds. Platypus TCR δ chain C region sequence (GenBank accession number XM_001516959) was used to identify the single copy platypus C δ on scaffold 588, which is separate from any of the scaffolds containing the putative platypus TCR μ sequences.

PCR and cDNA analyses

A spleen cDNA library constructed from tissue from a Tasmanian platypus was screened by PCR (9). All PCR primer sequences used in this study are presented in Table I. PCR amplification was performed using AdvantageTM-HF 2 PCR (BD Biosciences, Clontech Laboratories, Palo Alto, California) with the conditions: denaturation at 94 °C for 1 min for 1 cycle, followed by 34 cycles of 94°C for 30 s, annealing/extension at 62 °C for 4 min, and a final extension period of 68 °C for 5 min. Forward and reverse primers complementary to sequence internal to the platypus $C\mu$ exon were paired with primers in the $\lambda gt10$ vector used to construct the library to amplify clones containing the 5' and 3' un-translated regions (UTR) (10). This approach generated the partial cDNA sequences analyzed. Full-length platypus TCRµ cDNA sequences were isolated by PCR using primers complementary to 5' and 3' UTR. PCR products were cloned using TOPO TA cloning Kit (Invitrogen, Carsbad, CA) and sequenced using BigDye Terminator Cycle Sequencing Kit (Applied Biosystems, Foster City, CA). The GenBank accession numbers of the cDNA sequences described here are: clone 21, GU458338; clone 26, GU458339; clone 2.22, GU458341; clone 3815, GU475137; clone 1951, GU475138; clone 1953, GU475139; clone 1954, GU475140; clone 1955, GU475141; clone 4951, GU475142; clone 4942, GU475143; clone 786, GU475144; clone 6, GU458343; clone 17, GU475135; clone 2.34, GU458340; clone 10 GU264000; clone 36, GU475136; clone 4966, GU475145; clone 1.22, GU458342.

Phylogenetic analysis

Phylogenetic analyses were performed on nucleotide alignments using the MEGA4 program (11) with UPGMA (Unweighted Pair Group with Arithmetic mean), MP (Maximum parsimony), Neighbor-Joining (NJ) and ME (Minimum Evolution) methods. Amino acid translations were first aligned to establish gapping and then converted back to nucleotide using the BioEdit program (12).

The GenBank accession numbers of the sequences used in the phylogenetic analyses of TCRμ C and V region sequences were: Cβ sequences are Echidna, AY423735; Platypus, XM 001509180; Opossum, AY014507; Human, AF043178; Mouse, FJ188408. Cγ sequences are Opossum, DQ499632; Platypus, DQ011295; Human, X15019; Mouse, X03802. Cα sequences are Echidna, DQ011301; Platypus, XM 001507799; Opossum, AY014504; Human, FJ79357; Mouse, DQ186679. Cδ sequences are Platypus, XM 001516959; Human, M21624; Mouse, M37694; Bandicoot, AY955295; Opossum, XP 001379771; Wallaby, AY238447; Frog, GQ262033 and GQ262033; Chicken, XM 423780. Cμ sequences are Wallaby, AY956350; Bandicoot, AY955293; Opossum Cμ sequences are from MonDom5 scaffold 3.430000001-435000000 (13). The sequences of platypus Cµ used in the alignment are from platypus assembly version 5.0.1 and scaffold locations are presented in Supplementary Fig. 1. Wallaby Vδ48, AY238448; Wallaby Vδ51, AY238451; Bandicoot Vδ46, DQ076246; Cattle Vδ13, D16113; Human Vα96, Z14996; Human Vα34, AB360834; Human Vβ04, M27904; Mouse Vβ16 M15616; Cattle Vβ19 D90129; Rabbit Vβ19 D17419; Sheep Vβ11, AF030011; Human Vγ29, M13429; Mouse Vγ38, M13338; Cattle Vγ 88, U73188; Sheep Vγ98, Z12998; Platypus Vγ95, DQ011295; Platypus Vγ19, DQ011319; Shark NAR62, AY114762; Shark NAR78, AY114978; Shark NAR82, AY261682; Shark NAR60, EU213060; Shark TNAR05, DQ022705; Shark TNAR88, DQ022688; Shark TNAR10, DQ022710; Opossum Vμ sequences are DQ979402, DQ979398, EF503722, EF5037719, DQ979397, DQ979396, EF503721, EF503718. The sequences of platypus Vµ used in the alignment are from platypus assembly version 5.0.1 and locations are presented in Supplementary Fig. 1. Frog VHδ sequences are GQ262028, GQ262032, GQ232013; IgVH sequences are: possum VH50, AAL87470; possum VH1, AAL87474; bandicoot VH5.1, AY586158; Opossum VH sequences are from MonDom5 scaffold 1.295000001-300000000 (12). Mouse VH3660, K01569; mouse VH3609N, X55935; mouse VHDNA4, M20829; mouse VHJ588, Z37145; mouse VHJ606, X03398; mouseVHQ52, M27021; mouse VHS107, J00538; mouse VHSM7 M31285; mouse VH11, Y00743; mouse VH98, AJ851868. Human VH sequences were obtained from the VBASE database. Pig VH3, U15194; cattle VH, AF015505; sheep VH, Z49180; echidna VH7g, AY101438; echidna VH8g, AY101439; echidna VH51g, AY101442; platypus VH29, AF381294; platypus VH26, AF381293; platypus VH3, AF381314; platypus VH53, AF381304.

Results

Identification of a platypus TCRµ homologue

Fifteen gene sequences with similarity to opossum $C\mu$ were identified in the platypus whole genome assembly (14). Searching the unassembled, raw trace sequences from the platypus whole genome shotgun sequence did not uncover any additional genes with homology to opossum $C\mu$ Six of these contained complete open reading frames (ORF) and were used in all subsequent analyses (Supplementary Figs. 1 and 2). When compared to opossum $C\mu$ and conventional TCR C genes from a variety of mammals, the platypus sequences had greatest nucleotide identity to opossum $C\mu$ (Table II, Supplementary Fig. 2). Included in these analyses was the single copy conventional platypus TCR δ C gene which is located on scaffold 588 in the genome assembly, separate from any of the TCR μ related genes (Table II, Fig. 1 and data not shown). Phylogenetic analyses using several models for tree reconstruction result in the platypus and marsupial $C\mu$ together forming a well-supported monophyletic clade consistent with having identified the platypus TCR μ homologue (Fig. 1).

Platypus TCRµ is transcribed in a double V form

To investigate the structure of expressed platypus TCRµ, full-length transcripts were isolated from a spleen cDNA library. Transcripts averaged 1300 bp in length, which is longer than a conventional TCR transcript and more similar to the double V encoding opossum TCRµ (Fig. 2). Each encoded a leader (L) peptide followed by two complete V domains, designated V1 and V2 for the 5' (N-terminal) and 3' (C-proximal) domains, respectively. They also contained one C domain along with sequences corresponding to the connecting peptide (CP), transmembrane (TM) and cytoplasmic (CT) regions typical of trans-membrane TCR chains (Fig. 2). The clones encoded conserved residues found in conventional TCR including cysteines forming intra-chain disulfide bonds in the V and C domains as well as inter-chain disulfide bond in the CP (Fig. 2). The framework region (FR) 4 of V1 and V2 contain the sequence YGXG and FXXG, respectively, similar to the conserved FGXG motif in conventional TCR and marsupial TCRµ (4,15,16) (Fig. 2). Also present are two positively charged amino acids (arginine and lysine) in the TM region that, in conventional TCR chains, participate in association with the CD3 signaling complex (17). Comparison to the genomic sequence revealed that the CP is unusual in platypus TCRµ in that it is encoded on two exons, designated CP1 and CP2 with the conserved cysteine in CP2 (Fig. 2). This is unlike the opossum TCRμ and most conventional TCR where the CP is encoded by a single exon (4).

Both V1 and V2 are encoded by somatically recombined genes

The germ-line genes encoding the V1 and V2 domains were identified by comparing 18 unique V1 and 16 V2 sequences from both partial and full-length platypus splenic cDNA clones to the genome assembly. V1 and V2 domains share less than 65% nucleotide identity to each other and, by convention, are encoded by different V gene sub-groups designated $V\mu 1$ and $V\mu 2$, respectively. Nine $V\mu 1$ and six $V\mu 2$ genes were identified in the germ-line sequence (Supplementary Fig. 1). All nine of the Vµ1 genes contained upstream exons encoding a conserved L sequence; however none of the Vµ2 germ-line genes had a L exon (not shown). The sequences corresponding to FR4 in V1 and V2 were also used to identify eight Jμ1 and twelve Jμ2 genes, respectively. Jμ1 and Jμ2 are easily distinguished by length and sequence with Jµ1 being shorter and sharing less than 50% nucleotide identity with Jµ2 genes (Fig. 3). All Vµ and Jµ genes were flanked by conserved RSS, the recognition substrates for the Recombination Activating Gene product (18). The RSS flanking the V_µ and Jµ genes contained 23 and 12 bp spacers, respectively, typical of TCR genes (Fig. 3). In all cDNA sequences analyzed, $V\mu 1$ were recombined to $J\mu 1$ and $V\mu 2$ to $J\mu 2$. These results support both the V1 and V2 domains in platypus TCRµ are encoded by exons that are fragmented in the germ-line DNA and undergo RAG mediated V(D)J recombination.

The sequences corresponding to complementarity determining region 3 (CDR3) differed both in length and diversity between V1 and V2 domains (Fig. 2). The V1 CDR3 are longer and up to 22 codons in length whereas none of the V2 CDR3 exceeded 12 codons. Using the V1 CDR3 sequences identified 35 putative D μ genes in the platypus genome assembly, all of which were asymmetrically flanked by RSS containing a 12 bp spacer on the 5' side and 23 bp spacer on the 3' side, as is typical of TCR D genes (Supplementary Fig. 3). Based on length and nucleotide identity the D genes fell into two groups designated D μ 1 and 2. D μ 1 (n = 20) contained coding regions 10 to 13 nucleotides in length while D μ 2 (n = 15) were 18 to 19 nucleotides (Supplementary Fig. 3). There was greater than 75% nucleotide identity within each group but less than 40% nucleotide identity between D μ 1 and D μ 2 genes. Although D μ genes could be distinguished in the genomic sequence, individual contributions to the V1 junctions were difficult to establish due to their similarity and short length. Nonetheless it was possible to determine that the V μ 1-J μ 1 junctions contained two, three or four D μ genes, in roughly a 1:2:1 ratio, similar to the multiple D genes found in

opossum TCR μ rearrangements (Fig. 4, Supplementary Table 1). Typical of D gene segments, the D μ present in V1 junctions were used in multiple reading frames (Supplementary Fig. 3). The gene segments encoding the V1 domains demonstrated extensive trimming and no evidence of P nucleotide additions, although N nucleotide additions were common (Fig. 4).

In contrast to V1, the CDR3 of 14 of the 16 V2 cDNA sequences could be accounted for entirely by recombination between germ-line V μ 2 and J μ 2 genes, with evidence for P and N nucleotide additions but no D μ genes being incorporated (Fig. 4, Supplementary Table 1). The remaining two clones contained a short stretch of four or five nucleotides that matches D μ 2.8, and cannot be ruled out as being from a D segment. Whether coincidence or evidence of a D segment is not clear, and is not evident from the genomics where no D μ has been found between V μ 2 and J μ 2 gene segments (see below). These results are consistent with the longer CDR3 in V1 domains being due to incorporation of multiple D segments and the shorter V2 CDR3 being the result of direct V to J recombination in most if not all junctions.

Platypus TCRµ V genes are related to clan III VH genes

The relationship $V\mu$ genes have to each other and with V genes from Ig and conventional TCR was investigated by phylogenetic analyses. These analyses included VH from the platypus IgH locus (19). The results of these analyses support $V\mu1$ and $V\mu2$ each forming their own distinct clades with strong bootstrap support (99-100%) consistent with their designation as separate subgroups (Fig. 5). Furthermore, the platypus $V\mu$ subgroups together form a single clade nested within mammalian clan III VH genes. This is in contrast to the marsupial $V\mu$ ($V\mu$ and $V\mu$ j), which are not monophyletic but are closely related to VH (Fig. 5) (4).

Platypus TCRµ genomic organization

The $TCR\mu$ locus is not fully assembled in the current version of the platypus genome, but rather was scattered on 55 separate scaffolds ranging in length from less than 1 kb up to 64.8 kb (Supplementary Fig. 1). Seventeen of the 35 Dµ segments were on scaffolds also containing Vµ, Jµ and/or Cµ sequences, supporting their being part of a larger $TCR\mu$ locus (Supplementary Fig. 1). Combining the scaffold analyses with the cDNA sequences reveals a minimal model for the organization of the platypus $TCR\mu$ locus. Three scaffolds contain multiple Dµ either transcriptionally downstream of Vµ1 genes (scaffold 3930) or upstream of a Jµ1 gene (scaffolds Ultra190 and 19044) consistent with the evidence from cDNA sequences having multiple Dµ in the junctions between Vµ1 and Jµ1 genes (Fig. 6A, Supplementary Fig. 1). One scaffold (28416) contains single Vµ2 and Jµ2 genes that correspond to those used in expressed recombinations (Fig. 6A, Supplementary Fig. 1, Supplementary Table 1). However, no Dµ genes were found on this scaffold consistent with the lack of D segments in the majority of Vµ2-Jµ2 junctions (Figs. 4 and 6, Supplementary Fig. 1, Supplementary Table 1).

Full-length cDNA clones containing similar or identical $V\mu 1$ sequence also had similar or identical $J\mu 1$, $V\mu 2$, $J\mu 2$ and $C\mu$ (Supplementary Table 1). The most parsimonious explanation for these observations is a cluster organization of platypus $TCR\mu$ genes, similar to that found in marsupials (4). In other words, the V, D and J genes encoding V1 domains are upstream of the V and J gene segments encoding V2, followed by $C\mu$ (Fig. 6B). Consistent with this prediction, three scaffolds (19044, 26255, and 33931) contain $J\mu 1$ genes upstream of $V\mu 2$ genes and many of the scaffolds containing $C\mu$ genes also contained an upstream $J\mu 2$ (Fig. 6A, Supplementary Fig. 1). A conservative model for the organization of the platypus $TCR\mu$ genes is presented in Figure 6B. The model may be overly conservative

since two cDNA clones appeared to use different $V\mu 1$ but the same $J\mu 1$ while two others appeared to use the same $V\mu 1$ recombined to two different $J\mu 1$ (compare clones 2.34, 10 and 17 in Supplementary Table 1). These results imply there may be multiple $V\mu 1$ and $J\mu 1$ in some clusters, or alternatively may be due to trans-cluster recombination as has been found for both opossum $TCR\mu$ and shark $TCR\delta$ genes (4, 20).

To estimate the possible number of TCR μ clusters, the number of unique C μ sequences that could be isolated from an individual platypus was determined. PCR was performed on genomic DNA from a single platypus using primers designed to amplify all 15 C μ identified in the genome assembly. Twenty individual clones were sequenced and yielded nine distinct C μ sequences consistent with at least five C μ exons per haploid platypus genome (not shown). This number is slightly lower but not significantly different from what would be predicted from the platypus whole genome sequence where 15 different C μ were identified or a minimum of eight per haploid genome. Whether this is an artifact of the assembly or normal platypus variation remains to be determined.

Discussion

The discovery of a platypus TCR μ homologue confirms that this unconventional TCR locus is not unique to marsupials but rather it is ancient in the mammalian lineage and appeared prior to the divergence of the prototherian (monotremes) and therian (marsupial and placental) mammals more than 165 MYA (7). TCR μ was clearly retained in the marsupial lineage and, therefore, would have been present in the last common ancestor of marsupials and placental mammals. However, no TCR μ homologue has been identified in placental mammals, consistent with gene loss in this lineage (2). Furthermore, a TCR μ homologue has yet to be found in the available avian, reptilian, and amphibian genomes, consistent with its appearance in the synapsids (mammals and their extinct relatives) after their divergence from the diapsids (birds and reptiles) 310 MYA (2, 21). This conclusion is also consistent with phylogenetic analyses of TCR μ C region genes published previously, where marsupial C μ appears to diverge from C δ after the split between mammals and birds (4).

The most distinctive feature common to both marsupial and platypus $TCR\mu$ is their transcription in a form predicted to encode three extra-cellular Ig domains (V-V-C) instead of the conventional two domains (V-C). TCR with this characteristic have only been described in one other vertebrate lineage, the cartilaginous fish. Both the elasmobranchs (sharks, rays, and skates) and the holocephalins (ratfish) use an isoform of $TCR\delta$, called NAR-TCR, that also has a double V expressed with a conventional $C\delta$ (22).

There are a number of common characteristics shared between mammalian TCR μ and shark NAR-TCR, as well as distinctive differences (Table III). In both platypus TCR μ and NAR-TCR the exons encoding both V domains require somatic DNA recombination to be assembled (22). The supporting or V2 domains in NAR-TCR are encoded by a dedicated subset of V δ gene segments that, like the platypus V μ 2, lack L sequences and would be unable to encode the N-terminus of an extra-cellular protein (22). This is different, however, in marsupials where the exon encoding the V2 domain, called V μ j, is pre-assembled as a germ-line joined gene and contains a L sequence that is contiguous with the exon encoding the extra-cellular V domain (Fig. 6C) (4). In the case of marsupial TCR μ this L sequence is left out of the V μ j exon in the mature mRNA due to a canonical RNA splice site at the junctions between the L and V sequences (2, 4). This arrangement makes it possible to transcribe a two-domain form of marsupial TCR μ that contains only V μ j and C region. Indeed, such transcripts are found in the opossum thymus, however, are rare in peripheral lymphoid tissues, leading to the current working hypothesis that it is the double-V form that is the mature, functional chain (4). Furthermore, in the opossum, *Monodelphis domestica*,

there are eight tandem clusters of $TCR\mu$ genes and in six of these the $V\mu$ j L sequences contain mutations rendering them non-functional (2, 4). Therefore, while the shark and platypus have fully deleted the L sequence of the supporting V, the L sequences in marsupials are apparently degenerating due to lack of use.

Both TCR μ and NAR-TCR utilize V domains more similar to antibody V genes than conventional TCR V genes. The N-terminal V domains in NAR-TCR are related to V used in IgNAR, which are light-chainless antibodies unique to cartilaginous fishes (22, 25). As already described, the second V in NAR-TCR is a V δ gene, making the NAR-TCR appear to be a hybrid between IgNAR and TCR δ (22). In contrast, the genes used to encode both V1 and V2 domains in platypus TCR μ are indistinguishable from mammalian clan III Ig VH genes and unrelated to NAR V genes. Marsupial V μ and V μ on the other hand are somewhat intermediary. V μ j are more similar to Ig VH, but do not fall within the three traditional mammalian VH clans, and V μ appears to be more related to NAR V genes, although this latter relationship is only weakly supported in phylogenetic analyses (Fig. 5).

The current model for the structure of NAR-TCR is an unpaired N-terminal domain, much like the V-NAR domain in IgNAR, binding antigens as a single domain (22, 25). This Ag binding is similar to that which has been described for single V domain IgNAR antibodies in sharks and light-chainless IgG in camels (26, 27). It seems likely that TCR μ is structured similarly to NAR-TCR, with a single, unpaired N-terminal V domain capable of binding antigen directly. Based on conserved residues, including cysteines TCR μ is predicted to form a heterodimer with another TCR chain (4). However, since no other TCR related genes encoding a three-domain chain have been found in the marsupial genome it is predicted that the partner is a conventional two domain TCR chain, likely TCR γ , leaving the N-terminal domain unpaired (2).

The common characteristics found in mammalian $TCR\mu$ and shark NAR-TCR raise the question of whether these features are due to homology by descent or convergent evolution. An argument could be made that the evolutionary distance between sharks and mammals is sufficiently vast, and the differences between $TCR\mu$ and NAR-TCR extensive enough that each evolved independently and appear analogous due to convergence on a common structure and function. This could imply a common evolutionary pressure shared between cartilaginous fish and early mammals to have T cells capable of binding Ag directly using single domain binding sites.

Phylogenetic analyses of platypus and marsupial TCRµ C region support that they are orthologous genes that would have been found in a last common ancestor of the three living mammalian lineages. However, following the divergence of the oviparous monotremes from the viviparous marsupials and placental mammals, TCRµ appears to have followed different evolutionary paths. In the placental mammals it was lost altogether (2). As discussed earlier, in the marsupials the genes encoding the V2 domain appear to have been replaced in the germ-line by a pre-joined V gene, most likely via retro-transposition (4). This novel marsupial adaptation is consistent with the V2 domains serving strictly supporting roles rather than being Ag-binding and, therefore, requiring little or no clonal variation. In the platypus the TCRµ V2 domain is encoded by somatically recombined genes, but variation remains restricted through limited junctional diversity, with no D segments and few N or P additions in the V-J junctions. Comparisons of the length of the CDR3 region in the platypus and marsupial V2 domains, where they are both relatively short, suggests that D segments, if they were ever present, were deleted early in the evolution of TCRµ prior to the divergence of prototherians and therians (4). The mean codon length of the platypus V2 CDR3 is the same (n = 11) as that found in the germ-line joined marsupial $V\mu j$ genes (Table III). In contrast the V1 domains of both platypus and opossum TCRµ have comparatively longer

and more diverse CDR3 due to the incorporation of multiple D segments during V(D)J recombination in both species (4).

The lack of an intron separating the L from the V in the V μ j exon is evidence of retro-transposition in the evolution of TCR μ in marsupials (4). In other words, V μ j is a functional, partially processed gene. The insertion of joined V genes into the germ-line by retro-transposition would require co-existing retro-elements in the genome and one noteworthy distinction between the opossum and the platypus genomes is the abundance of retro-elements. The opossum has among the highest percentage of retro-elements of any vertebrate genome sequenced (28). In contrast, monotremes are relatively devoid of retro-elements (14, 29). Whether this extreme difference contributed to the evolution of opossum and platypus TCR μ is not known. Furthermore, this explanation is not fully satisfying since processed pseudogenes have been found in the platypus and echidna genomes, consistent with retro-transposition having occurred sometime in the past for some monotreme genes (10).

Phylogenetic analyses support TCRμ being related to and likely derived from a TCRδ ancestor (4, 5). As stated earlier, if $TCR\mu$ evolved from a duplication of $TCR\delta$ genes it likely occurred after the separation of mammals from birds and reptiles (4). However, some insight into the origins of TCRµ may come from recent work on the genetics of amphibian TCR δ chains (23). The $TCR\alpha/\delta$ locus in the frog *Xenopus tropicalis* contains two C δ genes, one of which, $C\delta 1$, is expressed with V genes called VH δ . These frog VH δ that are indistinguishable from clan II Ig VH genes and, although the X. tropicalis $TCR\alpha/\delta$ and Igh loci are closely linked, the VH δ genes appear to be dedicated for use in TCR δ chains and are not used in IgH chains (23). This close linkage, however, may have facilitated insertion of VH genes among the TCR δ genes in amphibians. The region of the frog $TCR\alpha/\delta$ locus containing $C\delta 1$ and multiple VH δ genes is distinct and in an inverted transcriptional orientation from the rest of the $TCR\alpha/\delta$ genes, functioning almost as a separate mini-cluster (23). Amphibians, therefore, appear to be another vertebrate lineage that uses $TCR\delta$ chains containing antibody-like V genes. Unlike TCRμ and NAR-TCR, frog TCRδ chains are not expressed with two V domains, however. Rather, X. tropicalis TCRδ chains using VHδ are structured like conventional two-domain TCR chains.

It is possible, and seems likely, that the TCR μ locus evolved from genome duplication and translocation of an ancestral region of the $TCR\alpha/\delta$ locus similar to the C δ 1 region in frogs. Indeed, the discovery of VH genes in the *X. tropicalis TCRa/\delta* locus is consistent with their presence in the $TCR\delta$ locus prior to the evolution of TCR μ . Internal duplications of clusters of V, D, and J segments within the $TCR\mu$ locus, as hypothesized previously, would then give rise to the double V organization in mammals (2). What remains puzzling is the variation in the source of VH genes used in each lineage. The VH δ in *X. tropicalis* are apparently derived from clan II VH, the platypus V μ genes are clan III VH, and although the marsupial V μ genes are more closely related to VH than TCR V genes but fall outside the clan I, II, and III designation. These observations suggest that the V genes used in TCR δ or TCR μ chains have been replaced over time with different VH lineages, even within the mammals. If the platypus $TCR\mu$ locus is indeed organized as tandem clusters similar to what has been shown in opossum (4), such gene clusters may facilitate gene replacement and duplication that is not easily achieved by the translocon organization of the conventional TCR genes.

The lack of TCR μ in commonly studied mammals such as humans and mice no doubt contributed to it remaining undiscovered for nearly a quarter century following that of the conventional TCR α , β , γ , and δ (4, 28-31). Determining why placental mammals may have lost this TCR chain will require first determining what function(s) TCR μ ⁺ T cells perform in those species where they are found.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Special abbreviations

RSS recombination signal sequences

ORF open reading frame

NAR New Antigen Receptor

MYA million years ago

VH Ig heavy chain V region

L leader

CP connecting peptide

TM transmembrane

CT cytoplasmic

FR framework region

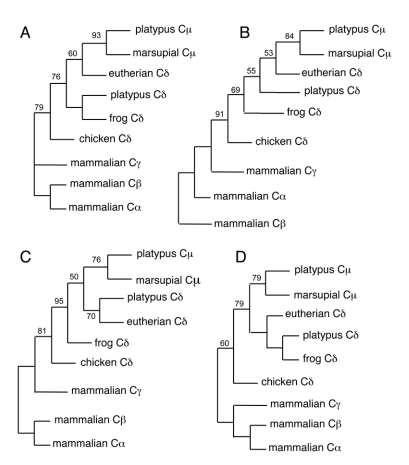


FIGURE 1.

Phylogenetic analyses of platypus and marsupial $C\mu$ and C regions from conventional TCR chains. Phylogenetic relationship between $C\mu$ and other conventional TCRs are simplified according to the phylogenetic trees constructed using different methods: (A) Neighbor-Joining (NJ); (B) Maximum Parsimony (MP); (C) Unweighted Pair Group Method with Arithmetic mean (UPGMA); (D) Minimum Evolution (ME). All phylogenetic analyses are based on nucleotide alignments and branch support is indicated as the percentage of out of 1000 bootstrap replicates.

	V1
21 26 2.34 2.22 1.22 6	0.4 Leader >< FR1 > CDR3 >< FR4 > CDR3 > FR4 > FR3 ST
	V2
21 26 2.34 2.22 1.22 6	FR1
	c
21 26 2.34 2.22 1.22 6	<pre></pre>
21 26 2.34 2.22 1.22 6	<pre>< CP1</pre>

FIGURE 2.

Predicted amino acid alignment of full-length platypus TCR μ cDNA clones. Dashes indicate identity and gaps introduced to the alignment are shown as dots. The sequences were divided into the Leader, V1, V2 and C domains. The FR and CDR of the V domains along with the C μ , CP, and TM-CT of C domain were shown above the sequence alignment. Conserved cysteines are shaded. Conserved lysines and arginines are shaded and indicated by *. Conserved residues YGXG and FXXG in FR4 of the V1 and V2 domains, respectively are noted. The borders of CDR and FR were indicated above the sequences.

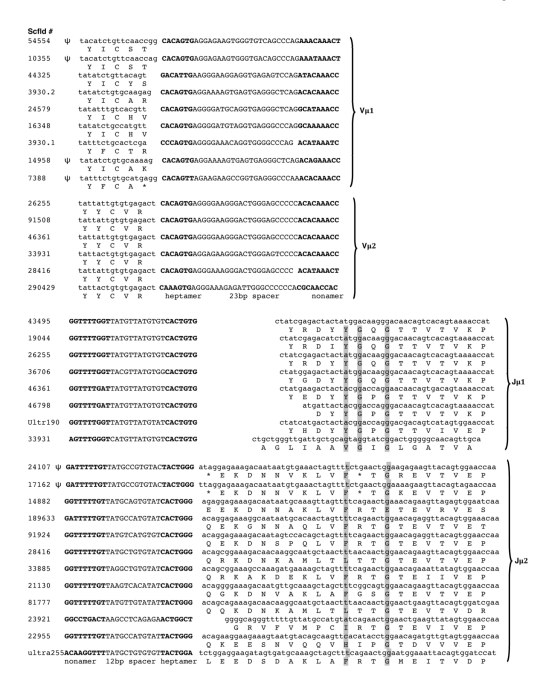


FIGURE 3.

Nucleotide sequence and translations of the 3' end of $V\mu 1$ and $V\mu 2$ gene segments and complete $J\mu 1$ and $J\mu 2$ gene segments. RSS flanking V and J gene segments in platypus genome are indicated. The scaffolds on which V and J sequences were identified are shown on the left. Pseudogenes are indicated by ψ . Stop codons are indicated as *. Nucleotide sequences of V and J genes are shown in lowercase with amino acid sequences underneath, whereas the RSS sequences are shown in uppercase. Heptamers and nonamers are in bold, and 12 bp or 23 bp spacers are indicated. The YGXG and FXXG conserved motif corresponding to FR4 are shaded.

Α													
Clones	No. V1 3' end	N		D		N	D	N	D	N	D	N	J1 5' end
21	CGGTACATCTGCCACGTT		A	AGGAGG		TCCAACGTCGGC	ACTGGA		ATAGG	TTGGG	GGACAA	С	TATGAAGACTACTATGGACCAGGAACAAGA
26	CGGTATATCTGCCACGTT	ATA	TCGCT	ACTAC		TCCCTTGGGGTT	TGGAAC	G	TGGG		GGAAC		GTGGGAACTACTACGGACCAGGAACAACA
2.22 3815	CGGTATATCTGCCACGTT	GG	GGG	AGGATGO AGACTG		AAGTTGATTTTC	TGGG	TTATGA	GGAAC	CTCCT	ATGCCT	CGCCCC	TATGAAGACTACTACGGACCAGGAACACCA
3815 1915	CGTTATATCTGCCACGTT		GTT	TGGAGC		AAGTTGATTTTC	GGAAC	GGGT	TGGG	GCCAT			TATGAAAACTACTACGGACCAGGAACACG
1915	CGGTATATCTGCCAC	ACAC	TICTIT	TGGAGC		ATATOTO	AGACTGGT	GGGT	TEGG	ATCCTATTAT	GAGGA		TACTACGGACCAGGAGCAACA
1954	CGGTATGTCTGTCACGTT		ng .	TGGAGC		TTGG	CTTA	GTTAT	GAGGA	C	GAGGA		TATGAGGACTACTACGGACCAGGAACAACA
1955	CGGTATATCTGT		TCGA	AGAGGAG		CT	ACTGGA	UTIAL	AGGGGG	CTGGT	TGGG	TCTTGGAGACTC	TATGAAAACCACTACGGACCAGGAACAACA
4951	CGGTATATCTGCCACGTT		CGT	AATATA		TGTCA	ACTGGT		TGGGC	TCTCT	TGGAT	CT	GAAAACTACTACGGACCAGGAACAACA
4942	CGTTATATGTGCCAC		GTATATTTAAAC	TGGCC		GTCTTTCGCTGGT	ATGAC						AGGAACAACA
786	GTATATCTGCCACGTC	GTCA	ATTATA	TATAGGAGAG	GAC	AGGGCT	ATGAC	ACC					CTACTACGGACCAGGAACAACA
6	CGGTACATCTGCCACGTT		ATATCATA	TTGG		CT	GGTC	GTTCTAGGTATA					ACTACTACGGACCAGGAACAACA
17	GATATATCTGCCACG		TCTTCGGT	TGGG		GCGCCGT	GTCAGGTT		TGGG	G			ACTACTACGGACGAGGAACAACA
2.34	CGCTACATTTGCCAC		GT	TGGG		GGT	TGGGC	ACCTCTG	GGAAC	AATCGGACC			CTATGAAAACCACTACGGACGAGGAACAACA
10	CGGTATATCTGTCACGTC		TACAGGA	ACTGGA		ATGGCGTGGGAG	AGGAGA	ACGAGGTTCAATCACAT	GGATGGAG	ACTAACACAT			GACCAGGGACGACAGTCAGAGTGCGACCAT
36	CGGTATATCTGT	GC	AAAA	TGGG		TTAAGAAGACGGT	GAAC	TGGCAATACCTGG	GGTC	CTATATTGACGTC			TACGGACCAGGGACAACC
4966 1.22	CGATACATCTGTGCAAGA CGGTACATCTGTGC	C	AGG	ATGACTGG		TC GTCAATATTATTTTGGCGGC	CTTA	TGCT					ACTACTACGGACCGGGGACTACAGT GACTACTACGGACCAGGGACAACA
A166	COGTACATCIGIGC			UGICAGE		GICAATATTATTTTGGCGGC							GRUTACTACOGRICAGOGRICAGUA
В													
Clones		P	N	D	P		2 5' end						
21	GGCCGCTATTATTGTGTGAGAC	T	TG			AGGCGAAAGACAGTAGT							
26	GGCCGGTATTATTGTGTG	C	GCCC	GATTA	C	GGAGACAGACAATAGT							
2.22	GGCCGGTATTATTGTGTGAGAC					ACAGGAGAAAGACAATAGT							
3815	GGCCGGTATTACTGTGTG	_ c	GCAC	GATTAG	c	GGAGACAGACAACAGT							
1951	GGCCGCTATTATTGTGTGAGAC		UA		C	GAAAGACAATAGTO							
1953 1954	GGCCGGTATTATTGTGTGAGAC					ACGGAGAAAGACAATAGT							
1954	GGCCGGTATTATTGTGTGCGAC	.1	GGA			ACAGGAGAAAGACAATAGT ACAGGAGGAAGACAATACT							
1955 4942	GGCCGGTATTATTGTGTGAG		GGG			ACAGGAGGAAGACAATACT							
4951	GGCCGGTATTATTGTGTGCGAC		000			ACAGGAGAAAGACAGTACT							
6	GGCCGGTATTATTGTGTGAGAC					ACAGGAGAGAGACAATAGT							
17	GGCCGGTATTATTGTGTG		GA		GT	ACAGGATAAAGACAATAGT							
4966	GGCCGGTACTACTGTGTGAGAG		U.A.		-	ACAGGGGGGGGGACAATGTT							
1,22													
		T AG					GCAAAGCTAGCTT	TOGGCAGT					
2.34	GGCCGGTATTACTGTGTGAGAG		TG		TG	AGGGAGAGACAACACCACCACCACCACCACCACCACCACC							

FIGURE 4.

Sequences corresponding to the CDR3 of (A) V1 and (B) V2 domains from full-length and partial platypus splenic TCR μ cDNAs.

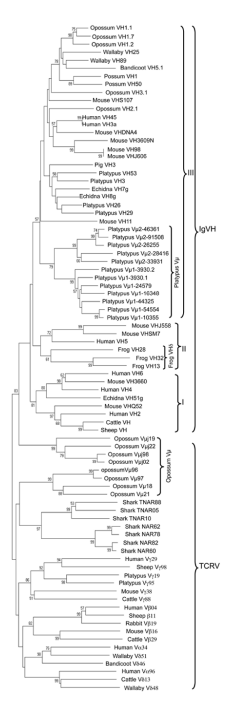


FIGURE 5.

Phylogenetic analysis of platypus and marsupial $V\mu$ including V genes from conventional TCR, shark NAR and NAR-TCR and Ig VH. This neighbor-joining tree is based on nucleotide alignments and branch support is indicated as the percentage of out of 1000 bootstrap replicates. Only those nodes with greater than 50% support are indicated. The three major clans of vertebrate VH are indicated by Roman numerals.

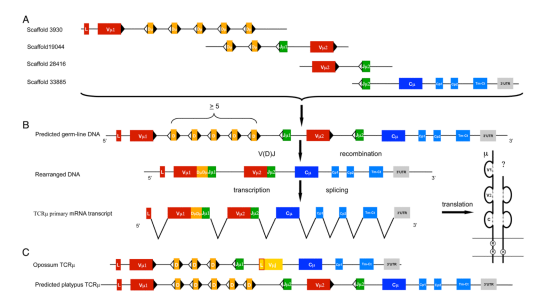


FIGURE 6.

Diagrams of the predicted platypus TCR μ gene organization, transcripts, and protein structure. (A) Representative TCR μ scaffolds containing TCR μ coding sequences. Closed or open triangles flanking the V μ , D μ and J μ gene segments indicate the presence of 23- or 12-bp spacer RSS, respectively. The L sequence, CP, TM-CT, and 3' UTR exons are indicated. (B) Predicted TCR μ germ-line DNA and rearranged DNA structure. (C) Primary TCR μ mRNA transcript structure. (D) Predicted TCR μ protein organization. Conserved R and K residues in the TM region are indicated. (E) Predicted platypus TCR μ cluster and representative opossum TCR μ cluster.

Table ISequences and description of oligonucleotide primers used

Sequence (5'-3')	Orientation	Region
CCTGGGCAGTGGGGGCCATGGCCTG	R	Сμ
GGGATAGTAATCTTTCACCAGGCAAG	R	Сμ
AGCAAGTTCAGCCTGGTTAAG	F/R	λ gt10 vector
ATTATGAGTATTTCTTCCAGGGTA	F/R	λ gt10 vector
CCCAACCCATGGTCTTTGTCATG	F	Сμ
GGAACCAGAGCTTCGCTGCTTGCC	F	Сμ
AACCATGCTGGTCCAGGTC	F	5' UTR
CAGGAGGGAAATGATTCAGG	R	3' UTR
CGGAAACAAAGAAGGCAGA	R	3' UTR
CGTGAAATACTCGGGGGAAT	F	$V\mu 1$
AGGCTCTGCATTGATCTTCG	F	Vμ2

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Table II

Comparison of platypus $C\mu$ with opossum $C\mu$ and conventional mammalian TCR C regions

	Platypus Cμ N=6	Platypus Cô N=1	Platypus C μ N=6 Platypus C δ N=1 Opossum C μ N=8 C δ^4 N=5 C α^5 N=5 C β^6 N=5 C γ^7 N=4	C8 ⁴ N=5	$C\alpha^5$ N=5	$C\beta^6$ N=5	$C\gamma^7 N=4$
Platypus Cμ N=6 ^I	80-98 ² (84 ³)	43-47(44)	50-56(52)	41-54(50)	41-54(50) 21-26(24) 25-32(29)	25-32(29)	27-33(31)
Platypus Cδ N=1	43-47(44)	100	43-47(45)	46-53(50)	26-30(28)	29-32(30)	29-33(32)
Opossum Cµ N=8	50-56(52)	43-47(45)	75-96(83)	41-54(48)	21-30(25)	26-34(30)	26-33(29)
C8 N=5	41-54(50)	46-53(50)	41-54(48)	55-83(67)	21-30(25)	24-31(27)	28-34(31)
$C\alpha N=5$	21-26(24)	26-30(28)	21-30(25)	21-30(25)	45-87(54)	22-33(27)	24-33(27)
Cβ N=5	25-32(29)	29-32(30)	26-34(30)	24-31(27)	22-33(27)	63-93(72)	28-36(31)
$C\gamma N=4$	27-33(31)	29-33(32)	26-33(29)	28-34(31)	24-33(27)	28-36(31)	48-76(54)

 $^{^{\}it I}$ Number of sequences included in the comparison

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The range of % nucleotide identity

 $^{^3}$ The mean % nucleotide identity

 $^{^{4}\}mathrm{C}\delta$ sequences of human, mouse, opossum, bandicoot, wallaby

 $^{5\}mbox{C}\alpha$ sequences of human, mouse, opossum, echidna, platypus

 $^{^6\}text{C}\beta$ sequences of human, mouse, opossum, echidna, platypus

 $^{^{7}}$ C γ sequences of human, mouse, opossum, platypus

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Comparison of the features of TCR μ , shark NAR TCR, and mammalian conventional TCR α/δ .

Locus	Model Species C gene Double V Somatic recombination	C gene	Double V	Somatic rec	combination	Leader sequence	edneuce	No. of D segments used	ments used	CDR3 leng	CDR3 length $^{\$}$ (mean)	V nature		Ref
				N-terminal	N-terminal C-proximal N-t	N-terminal	C-proximal	N-terminal	terminal C-proximal N-terminal C-proximal N-terminal C-proximal	N-terminal	C-proximal	N-terminal	C-proximal	
{	Platypus	СĒ	Yes	Yes	Yes	Yes	No	2-4	03	9-22(14)	9-12(11)	VH Clan III	VH Clan III herein	herein
ICK	Opossum	С	Yes	Yes	8 oN	Yes	$\mathrm{Yes}^{\#}$	1-3	NA*	8-29(17)	11	VH-related	VH-related	4
NAR-TCR	Nurse shark	CS	Yes	Yes	Yes	Yes	No	1	1-2	9-25(16)	9-27(16)	V-NAR	Vδ	22
TCRa/δ	Xenopus	C81	No	Yes	NA	Yes	NA	1-2	NA	7-20(13)	NA	$V\alpha,V\delta,VH\delta\;(VHClanII)$	NA	23
TCRa/8	Human	CS	No	Yes	NA	Yes	NA	2-3	NA	8-12(15)	NA	να, νδ	NA	24
TCRa/8	Mouse	CS	No	Yes	NA	Yes	NA	2	NA	6-19(13)	NA	$V\alpha, V\delta$	NA	24

\$ Range in codons

 § The C proximal V in marsupial TCRµ is a germ-line joined V

 $\slash\hspace{-0.4em}Fused$ to the V domain exon as the result of retrotrans position

* Not applicable